

REPORT

Genotype \times Adiposity Interaction Linkage Analyses Reveal a Locus on Chromosome 1 for Lipoprotein-Associated Phospholipase A₂, a Marker of Inflammation and Oxidative Stress

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Because obesity leads to a state of chronic, low-grade inflammation and oxidative stress, we hypothesized that the contribution of genes to variation in a biomarker of these two processes may be influenced by the degree of adiposity. We tested this hypothesis using samples from the San Antonio Family Heart Study that were assayed for activity of lipoprotein-associated phospholipase A₂ (Lp-PLA₂), a marker of inflammation and oxidative stress. Using an approach to model discrete genotype \times environment (G \times E) interaction, we assigned individuals to one of two discrete diagnostic states (or “adiposity environments”): nonobese or obese, according to criteria suggested by the World Health Organization. We found a genomewide maximum LOD of 3.39 at 153 cM on chromosome 1 for Lp-PLA₂. Significant G \times E interaction for Lp-PLA₂ at the genomewide maximum ($P = 1.16 \times 10^{-4}$) was also found. Microarray gene-expression data were analyzed within the 1-LOD interval of the linkage signal on chromosome 1. We found two transcripts—namely, for Fc gamma receptor IIA and heat-shock protein (70 kDa)—that were significantly associated with Lp-PLA₂ ($P < .001$ for both) and showed evidence of *cis*-regulation with nominal LOD scores of 2.75 and 13.82, respectively. It would seem that there is a significant genetic response to the adiposity environment in this marker of inflammation and oxidative stress. Additionally, we conclude that G \times E interaction analyses can improve our ability to identify and localize quantitative-trait loci.

From studies of humans and animal models, accumulating evidence has suggested a positive association between measures of adiposity and biomarkers of inflammation and oxidative stress.^{1–11} At the molecular level, this association seems to arise from the increased expression of adipokines in the white adipose tissue^{1,2} of central adipose depots.^{3–5} As a result of these advances, a unifying theory on the etiology of the metabolic syndrome posits that obesity leads to a state of chronic, low-grade inflammation and oxidative stress and that it is this pathological condition—of chronic inflammation and oxidative stress—that underlies most of the clinical sequelae associated with the metabolic syndrome.^{12–27}

On the basis of these findings, we hypothesized that the contribution of genes to variation in biomarkers of inflammation and oxidative stress may be influenced by the degree of adiposity—that is, “adiposity environment”—in individuals in whom they are expressed. We sought to test this hypothesis in the San Antonio Family Heart Study (SAFHS), which is a study of the genetic determinants of cardiovascular disease (CVD) in Mexican American families of San Antonio. We used the modeling approach of discrete genotype \times environment (G \times E) interaction, where two discrete adiposity environments—obese and nonobese—were defined according to criteria suggested by the World Health Organization (WHO).²⁸

The SAFHS population comprises large Mexican American extended families randomly ascertained with respect to CVD.²⁹ The SAFHS protocols were approved by the Institutional Review Board at the University of Texas Health Science Center at San Antonio, and all study participants provided written informed consent. The pedigree relationships exhibited by the sample population are reported in table 1.

Fasting blood samples were obtained from study participants at a clinic exam and were shipped the same day to Southwest Foundation for Biomedical Research (SFBR), San Antonio. Plasma and serum were isolated by low-speed centrifugation, and the buffy coat was harvested for DNA extraction.

We analyzed a biomarker of inflammation and oxidative stress in atherogenesis—namely, plasma activity of lipoprotein-associated phospholipase A₂ (Lp-PLA₂), which is also known as “platelet-activating factor acetylhydrolase” (PAF-AH).^{30–32} Plasma Lp-PLA₂ activity was determined using a commercial colorimetric assay (Cayman Chemical) with 2-thio-PAF as substrate and according to the manufacturer’s directions. Samples were run in duplicate, with average coefficients of variation of 2.5%. Enzyme activity was expressed in units of $\mu\text{mol}/\text{min}/\text{ml}$. Residuals from a least-squares multiple linear regression—with use of age, sex, age squared, oral-contraceptive use, and menopause

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Table 1. Pedigree Relationship Types in the SAFHS

Relationship	No. of Observed Pairs
Parent-offspring	2,550
Full siblings	1,780
Half siblings	260
Grandparent-grandchild	2,234
Avuncular	3,583
Half avuncular	498
First cousins	3,365

status as independent variables—were exactly normalized using an inverse Gaussian transformation in SOLAR (hereafter referred to as “rnLp-PLA₂”).³³ Anthropometrics, including height, weight, and waist and hip circumferences, were measured at a clinic exam as part of the SAFHS protocol. BMI, defined as the ratio of weight (kg) to height squared (m²), and waist-to-hip ratio (WHR), defined as the ratio of waist circumference to hip circumference, were determined from the anthropometric data.

It is by now well established that abdominal obesity is a major metabolic-syndrome risk factor.^{19,22,24,27} In an attempt to incorporate the abdominal-obesity component into a clinical definition of the metabolic syndrome, a WHO expert committee defined the abdominally obese as those with a combined BMI >30 kg × m⁻² and a WHR >0.90 in men and >0.85 in women.²⁸ We used these cutoffs to define a dichotomous adiposity environment, as given by the indicator variable

$$f_i = \begin{cases} 1 & \text{if obese} \\ 0 & \text{if nonobese} \end{cases}$$

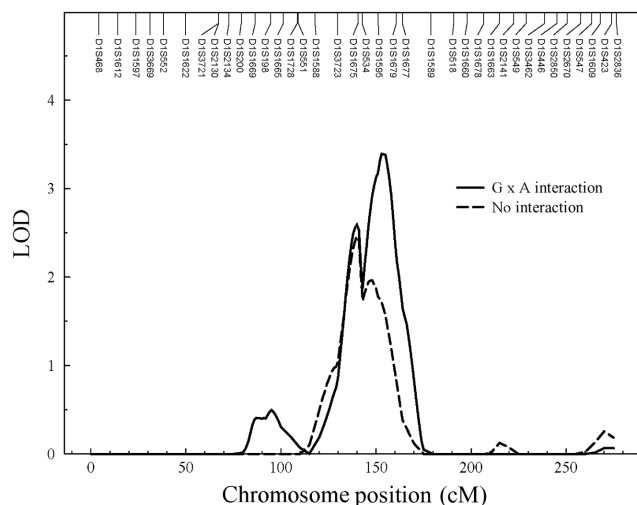
The sample size of individuals with data for rnLp-PLA₂, BMI, and WHR combined is 1,341.

DNA extracted from lymphocytes was used in PCRs for the amplification of individual DNA (*N* = 1,339) at 432 dinucleotide-repeat microsatellite loci (i.e., STRs), spaced ~10 cM apart across the 22 autosomes, with fluorescently labeled primers from the MapPairs Human Screening set, versions 6 and 8 (Research Genetics). PCRs were performed separately, according to manufacturer specifications, in Applied Biosystems 9700 thermocyclers. The products of separate PCRs, for each individual, were pooled using the Robbins Hydra-96 Microdispenser, and a labeled size standard was added to each pool. The pooled PCR products were loaded into an ABI PRISM 377 or 3100 Genetic Analyzer for laser-based automated genotyping. The STRs and standards were detected and quantified, and genotypes were scored using the Genotyper software package (Applied Biosystems).

Mistyping analyses were performed on the preliminary genotype-marker data with SimWalk2, following the recommendations of the program developers for accounting for mistyping error, by (1) blanking the errant called alleles, (2) recalling them conditional on the analysis (i.e.,

reassigning them under a different allele designation), or (3) retyping the mistyped marker or markers as resources permitted.^{34,35} Our overall rate of blanking mistyped markers was 1.37%. These mistyping analyses allow investigators to account for Mendelian errors and spurious double recombinants, both of which can severely reduce the power of a linkage analysis if not accounted for.³⁵ After addressing mistyping errors (by blanking, recalling, or retyping), these genotype data were then used to compute maximum-likelihood estimates of allele frequencies in SOLAR.³³ Empirical estimates of identity-by-descent (IBD) allele sharing at points throughout the genome for every relative pair were computed using the Loki package, which uses Markov chain–Monte Carlo methods.³⁶ The multi-point IBD estimates are required under our variance-components modeling approach (see below). The SimWalk2 and Loki programs both require chromosomal maps. We used the set of high-resolution chromosomal maps provided by the research group at deCODE genetics, which are included in Web table E in the work of Kong et al.³⁷

Total RNA was extracted from 1,000 lymphocyte samples by use of QIAGEN RNeasy 96 kits, and concentration was determined spectrophotometrically by use of a NanoDrop. Integrity of resuspended total RNA was determined by electrophoretic separation and subsequent laser-induced fluorescence detection by use of the RNA 6000 Nano Assay Chip Kit on the Bioanalyzer 2100 with the 2100 Expert software (Agilent Technologies). Antisense RNA (aRNA) was synthesized and purified using the Ambion MessageAmp II Amplification Kit, following the Illumina Sentrix Array Matrix 96-well expression protocol. Biotin-16-UTP (Roche)–labeled aRNA was hybridized to Illumina



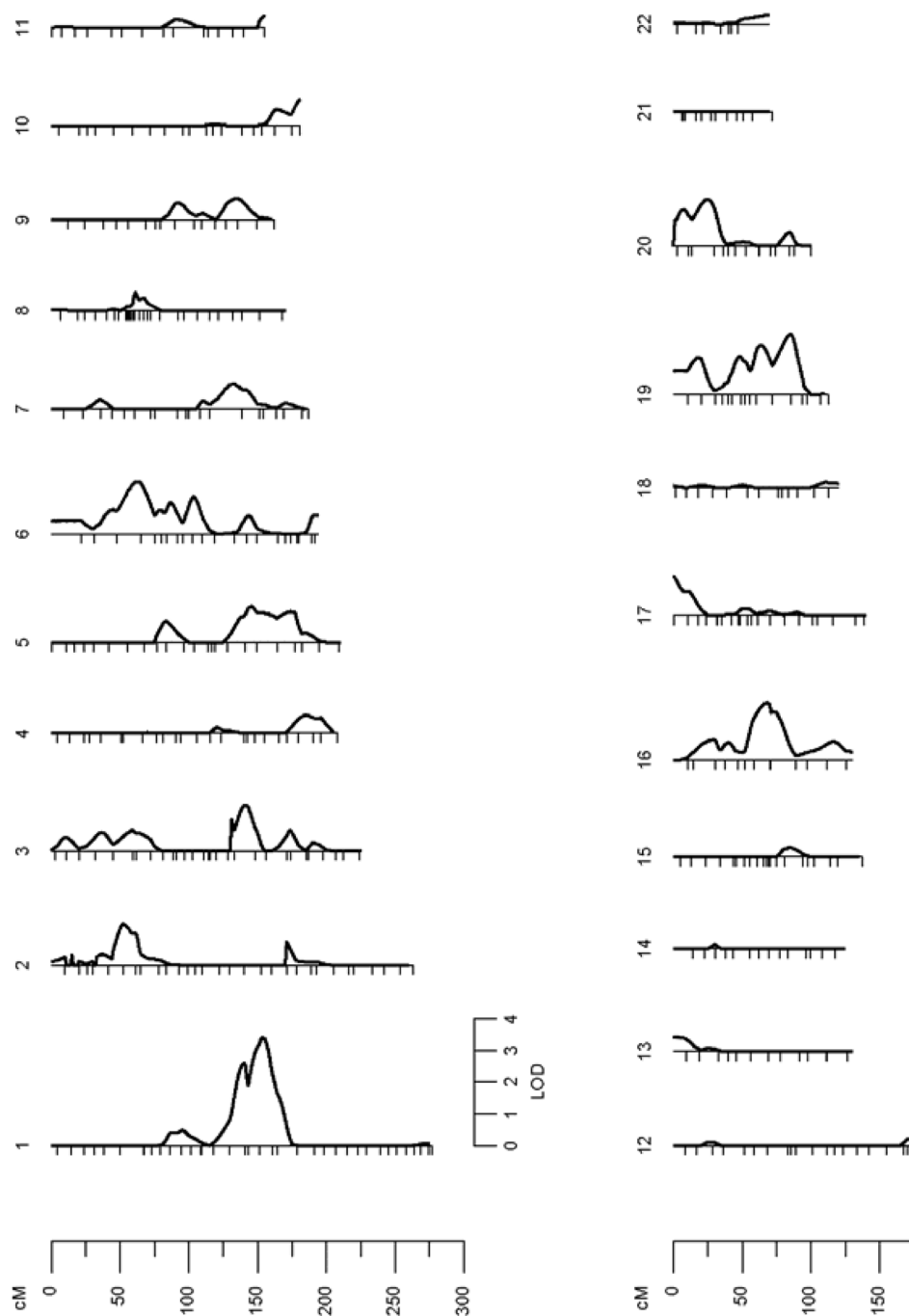


Figure 2. Multipoint genome scan of Lp-PLA₂

Sentrix Human Whole Genome (WG-6) Expression BeadChips. These BeadChips contain six arrays, each with 47,289 probes derived from human genes in the National Center for Bioinformatics Information (NCBI) Reference Sequence and UniGene databases. This system uses a “direct hybridization” assay, whereby gene-specific probes are used to detect labeled RNAs. Each bead in the array contains a 50-mer, sequence-specific oligonucleotide probe synthesized using Illumina’s Oligator in-house technology. Each array on a Human WG-6 BeadChip provides

genomewide transcriptional coverage of well-characterized genes, gene candidates, and splice variants. The Human WG-6 Expression BeadChips were scanned on the Illumina BeadArray 500GX Reader, a two-channel, 0.8- μ m-resolution confocal laser scanner, by use of Illumina BeadScan image data acquisition software (ver. 2.3.0.13). Illumina BeadStudio software (ver. 1.5.0.34) was used for data visualization and quality-control metrics.

To preclude confusion, we note that these data are expression levels of RNA transcripts and are not genotypic

data. They are treated herein as phenotypic data. It should also be noted that lymphocytes may not fully reflect the expression of all genes influencing adiposity, oxidative stress, and inflammation. However, there is growing use of lymphocytes as surrogate models for other tissues, such as neural tissues,^{38–40} and such work has generated substantial new discoveries.⁴⁰

The hypothesis of differential response to two environments is an example of the application of the theory of discrete $G \times E$ interaction.⁴¹ There is now a good number of published reports on the utility of this approach, both for understanding the relationship between genotype and environment in the process of phenotype determination and to aid in the identification and localization of QTLs.^{42–56} Under the theory of discrete $G \times E$ interaction, significant interaction arises for heterogeneity in the additive genetic variance (polygenic or QTL), an additive genetic correlation coefficient (polygenic or QTL) significantly different from unity, or both conditions.⁴¹ Therefore, to test for $G \times E$ interaction, we sought to falsify the null versions of the conditions that give rise to $G \times E$ interaction, which are a homogeneous additive genetic variance (polygenic or QTL) and/or an additive genetic correlation coefficient (polygenic or QTL) equal to 1.

For all possible pairwise combinations of values for the adiposity indicator variables f_x and f_z , where x and z are index individuals in the sample, the $G \times A$ interaction model covers three types of pairwise comparisons: within-obese, within-nonobese, and across-adiposity-environment comparisons:

$$\text{Cov}(y_x y_z) = \begin{cases} 2\phi_{xz}\sigma_{g0}^2 + \hat{\phi}_{xz}\sigma_{q0}^2 + \delta_{xz}\sigma_{e0}^2 ; \\ \quad \forall f_x = f_z = 1 \\ 2\phi_{xz}\sigma_{gn}^2 + \hat{\phi}_{xz}\sigma_{qn}^2 + \delta_{xz}\sigma_{en}^2 ; \\ \quad \forall f_x = f_z = 0 \\ 2\phi_{xz}\sigma_{g0}\sigma_{gn}\rho_{G(0,n)} + \hat{\phi}_{xz}\sigma_{q0}\sigma_{qn}\rho_{Q(0,n)} ; \\ \quad \forall f_x = 1, f_z = 0, \text{ or } f_x = 0, f_z = 1 \end{cases}, \quad (1)$$

where y is any given phenotype; $2\phi_{xz}$ gives the expected coefficient of relationship,

$$\phi_{xz} = \frac{1}{2}E\left[\frac{\kappa_{1j}}{2 + \kappa_{2j}}\right],$$

where the κ_{ij} are coefficients giving the j th locus-specific probability that a pair of relatives share i alleles IBD; $\hat{\phi}_{xz}$ is the estimated kinship coefficient based on marker data; δ_{xz} is defined as 1 when individuals x and z are the same and 0 otherwise; σ_{g0}^2 , σ_{gn}^2 , σ_{q0}^2 , σ_{qn}^2 , σ_{e0}^2 , and σ_{en}^2 are, respectively, the within-obese and within-nonobese additive polygenic, QTL, and environmental variances (the positive square roots of which give their corresponding SDs); and $\rho_{G(0,n)}$ and $\rho_{Q(0,n)}$ are the across-adiposity–environment additive polygenic and QTL correlation coefficients, respectively. We refer to this model as the “linkage interaction model.”

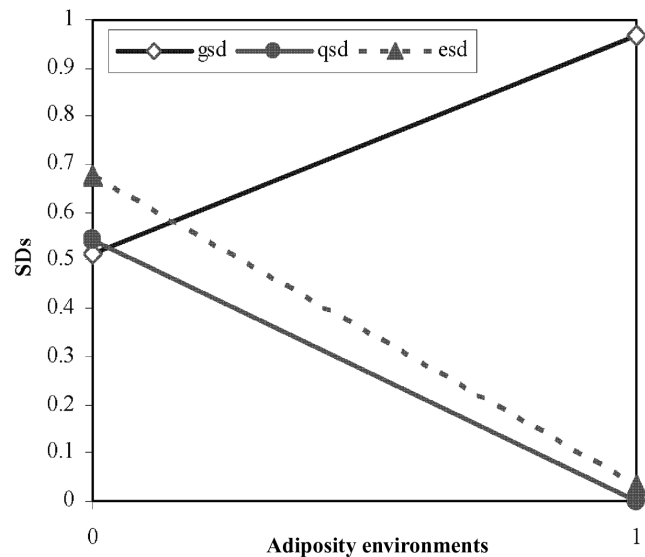


Figure 3. $G \times A$ interaction effects. The solid line with unblackened diamonds indicates genetic SD (gsd), the solid line with blackened circles indicates QTL SD (qsd), and the dashed line with blackened triangles indicates environmental SD (esd).

It will be necessary at this point to define the polygenic interaction model as a constrained version of the linkage interaction model in which the following constraint holds: $\sigma_{q0}^2 = \sigma_{qn}^2 = 0$.

The top and middle cases on the right side of the previous equation are the within-adiposity–environment versions of the standard linkage model used by Almasy and Blangero,³³ which hold for the obese and nonobese environments, respectively. The crucial part of the model is given by the bottommost case, which gives the covariance for the across-adiposity–environment comparison. Note that we are allowing for the possibility of heterogeneity in the residual environmental variance. This is necessary to preclude bias in detection of heterogeneity in the genetic-variance components. In all of our models, $\rho_{Q(0,n)}$ was constrained to equal 1, because the contribution to the model made by $\rho_{Q(0,n)}$ tends to be offset by the increase in degrees of freedom relative to the standard linkage model (results not shown).

We used SOLAR to perform genome screens under standard linkage and $G \times A$ interaction models across all 22 autosomes. For the standard linkage case, the likelihood ratio statistic, denoted by Λ , is distributed as $\frac{1}{2}\chi_0^2 + \frac{1}{2}\chi_1^2$.⁵⁷ It is important to note that SOLAR automatically corrects the LOD score for the standard case, to account for the above mixture distribution. The observed LOD scores under the $G \times A$ interaction model need to be further corrected because of the increase in degrees of freedom relative to the standard linkage model. Following Self and Liang,⁵⁷ it can be shown that when the polygenic interaction and linkage interaction models are compared, Λ is distributed as $\frac{1}{4}\chi_0^2 + \frac{1}{2}\chi_1^2 + \frac{1}{4}\chi_2^2$. We refer to this latter cor-

Table 2. Transcripts under the 1-LOD Interval of the Linkage Signal on Chromosome 1 that are Significantly Associated with Lp-PLA₂

Symbol	Name	Function	Location ^a (cM)	P ^b	Q
<i>CTMP</i>	C-terminal modulator protein	Protein kinase B regulation	147.18	.00174	.04653
<i>CTSS</i>	Cathepsin S	Elastase activity	147.64	5.15×10^{-4}	.01738
<i>SNX27</i>	Sorting nexin, family member 27	Intracellular sorting	148.52	4.81×10^{-4}	.01738
<i>FLJ23221</i>	Chromosome 1 ORF 54	ORF	148.78	.00164	.04653
<i>PBXIP1</i>	Pre-B-cell leukemia transcription factor interacting protein 1	Transcriptional regulation	151.85	2.77×10^{-4}	.01738
<i>SYT11</i>	Synaptotagmin, isoform 11	Mast-cell regulation	152.77	1.30×10^{-4}	.01738
<i>FCER1A</i>	Fc epsilon receptor 1A	Mast-cell activation	156.21	3.03×10^{-4}	.01738
<i>Hmm8932</i>	Hmm8932	Gnomon predicted gene	157.95	5.21×10^{-4}	.01738
<i>FCGR2A</i>	Fc gamma receptor 2A	C-reactive-protein receptor	158.42	1.52×10^{-4}	.01738
<i>HSPA6 = HSP70</i>	Heat-shock protein (70 kDa)	Chaperone-protein folding	158.44	4.18×10^{-4}	.01738

^a Locations are averaged interpolations against our map, with use of physical distances obtained from the University of California–Santa Cruz (UCSC) genome browser (Human [*Homo sapiens*] Genome Browser Gateway) for the two markers flanking the linkage peak at 153 cM.

^b P value of the beta coefficient for Lp-PLA₂ in a linear model in which the transcript is the dependent variable.

rection for increase in degrees of freedom as the corrected LOD score. Since the LOD score is equal to $\frac{\Lambda}{2\ln(10)}$, we can obtain a corrected LOD score on the basis of the appropriate distribution. To test the null hypothesis of homogeneity in the QTL variance (i.e., $\sigma_{qo}^2 = \sigma_{qn}^2$) at the genome-wide maximum (the point along the genome that has the highest LOD score), we performed likelihood-ratio tests. For model comparisons in which the QTL variances are constrained to be equal under the null hypothesis and in which the QTL variances are free to vary under the alternative hypothesis, Λ is distributed as χ_1^2 .

We found that rnLp-PLA₂ has a heritability of 0.55 ($P = 2.07 \times 10^{-41}$). Using the G \times A interaction model, we found a corrected, genomewide maximum LOD of 3.39 at 153 cM on chromosome 1 near marker *DIS1595* (figs. 1 and 2). A LOD score of 3.0 is taken as indicative of genomewide significance.⁵⁸ Moreover, using an approach based on the work of Feingold et al.⁵⁹ and implemented in Gauss 6.0.17 (Aptech Systems), we can compute the genomewide P value that corresponds to our LOD score of 3.39. This approach takes into account the finite marker density in the linkage map used in the multipoint QTL screens and the mean recombination rate for the pedigree population studied. The genomewide P value computed under said approach is .01477. In contrast, maximization of models lacking G \times A interaction did not provide strong evidence of a QTL anywhere in the genome. There is, however, a suggestive LOD score of 2.46 at 140 cM on chromosome 1 under the standard linkage model, with a corresponding genomewide P value of .14402. To highlight the improvement given by the incorporation of interaction effects, the results under the standard linkage model for chromosome 1 are also displayed. It will be noted that the location of the maximum LOD score changes from 140 cM on chromosome 1 under the standard linkage model to 153 cM on chromosome 1 under the G \times A interaction model. One plausible explanation is that the shift in peaks is simply the result of a model that affords a more precise signal location. In keeping with this view, Blangero et al. showed that G \times E interaction models increase the power to detect linkage signals and

precision of signal location.⁶⁰ Another plausible explanation is that there is another gene located at the linkage peak under the standard linkage model and that the G \times A interaction model recovers information that, in the aggregate, points to another location, harboring another gene, as the maximum while at the same time retaining the peak first observed under the standard linkage model. We cannot distinguish between these alternatives at the present time.

We found significant QTL G \times A interaction for rnLp-PLA₂ at the genomewide maximum on chromosome 1 ($P = 2.32 \times 10^{-4}$) (fig. 3). Specifically, the QTL additive genetic variance decreased from the nonobese to the obese environment. This finding may indicate a gene that is negatively regulated by adipose tissue. For instance, adiponectin is negatively associated with obesity,^{61,62} and a regulatory protein that both is related to the adiponectin gene and is itself negatively regulated by the proinflammatory cytokine, tumor necrosis factor- α (TNF- α), has been shown to be located in a region encompassing our linkage signal.^{63–65} Additionally, the gene for the adiponectin receptor AdipoR1 has been mapped to a broad region on chromosome 1 that encompasses our linkage signal.⁶⁶ Interestingly, the polygenic additive genetic variance exhibited significant heterogeneity (i.e., polygenic G \times A interaction) ($P = 3.54 \times 10^{-4}$) (fig. 3) and increased from the nonobese to the obese environment. This observation is consistent with the knowledge that adiposity promotes the expression of proinflammatory cytokines, such as TNF- α .^{12–27} Our findings of decreasing QTL variance and increasing polygenic variance in response to the adiposity environment may be reflective of down-regulated and up-regulated signals related to the inflammation response. The additive environmental variance also significantly decreased from the nonobese to the obese environment ($P = 6.86 \times 10^{-4}$) (fig. 3). This observation is consistent with the way the determinative system affecting Lp-PLA₂ comes under relatively more genetic control when going from the nonobese to obese environment.

To our knowledge, there are at least three other genome-scan studies that have reported signals on chromosome 1

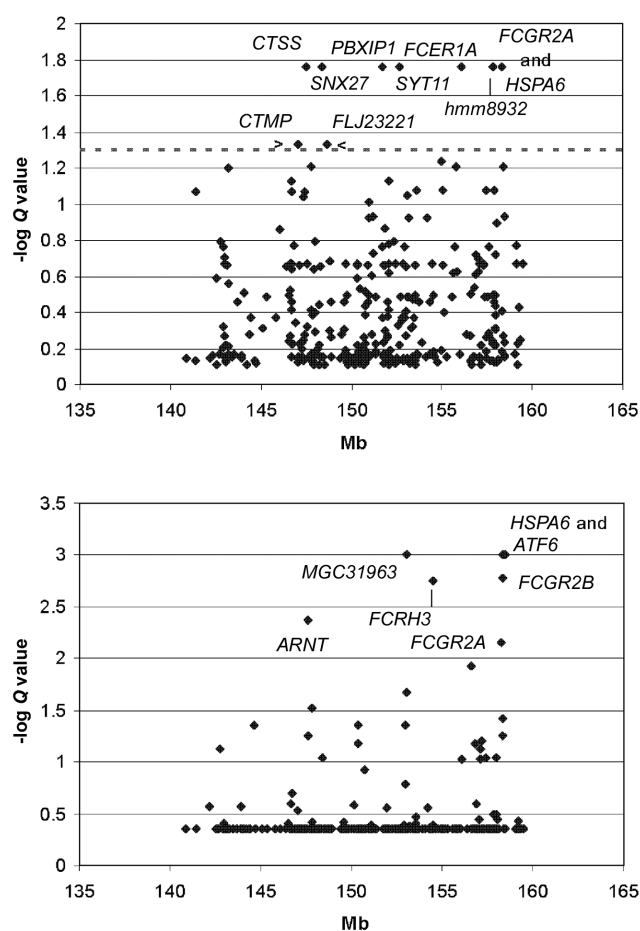


Figure 4. Transcripts under the linkage signal; transcripts within the 1-LOD interval of the linkage peak at 153 cM. *Top*, Negative logarithm (base 10) of the Q values for the test of association between the transcript and Lp-PLA₂ plotted against chromosomal location. The dashed line corresponds to the Q value threshold of .05 for the beta coefficients. *Bottom*, Negative logarithm (base 10) of the Q values computed from the pointwise LOD scores plotted against chromosomal location. The grid line at 2.0 corresponds to the Q value threshold of .01 for the pointwise LOD scores. Location is expressed in terms of megabases along the ordinate, for better separation. To plot all values on a scale at which the Q value threshold could be easily discerned, the three highest $-\log Q$ values for *MGC31963*, *HSP70*, and *ATF6* were arbitrarily given a $-\log Q$ value of 3.0. Their real corresponding Q values are in table 3 (*HSPA6* = *HSP70*).

in the vicinity of our signals for traits related to obesity and/or the metabolic syndrome.^{67–69} Ng et al.,⁶⁷ in the Hong Kong Family Diabetes Study, reported their genome-wide maximum LOD of 4.5 at chromosome 1q near marker *DIS1653* for the metabolic syndrome. Marker *DIS1653* is located at 151.68 cM on the deCODE map and is very near our genomewide maximum for the $G \times A$ interaction analyses. In the Framingham Heart Study,⁶⁸ Dupuis et al. reported a LOD of 3.86 at chromosome 1q near marker *DIS1679* for monocyte-chemoattractant pro-

tein-1 (MCP-1). Marker *DIS1679* is positioned at 160 cM in our chromosome 1 map, which is just at the outer margin of the 1-LOD interval in our $G \times A$ interaction analyses. Moreover, similar to Lp-PLA₂, MCP-1 is another biomarker of vascular inflammation. In a study of nuclear families ascertained at the University of Pennsylvania,⁶⁹ Reed et al. reported a LOD equivalent of 2.2 on chromosome 1 near marker *DIS484* for plasma cholesterol levels. Marker *DIS484* is at 157.51 cM on the deCODE map, which again is within the 1-LOD interval in our $G \times A$ interaction analyses.

We analyzed the gene-expression data to further characterize the area under the linkage peak. Because there are exactly 341 transcripts within the 1-LOD interval around the maximum, we employed a statistical methodology, based on the false-discovery rate (FDR) and Q Value concepts, to deal with the multiple-testing problem.^{70–74} The literature on the detection of differential gene expression in DNA microarray data—and similar such cases of data mining in statistical genetics and genomics—seems to point to methods that control the FDR rather than to the highly conservative method of controlling for the familywise error rate, such as the well-known Bonferroni correction.^{70–81} The Q value is mathematically defined as the minimum positive FDR (pFDR) observed for a set of significant results: $Q \text{ value} = \min \{pFDR\}$. It is a measure of the proportion of false-positive results expected on declaring a particular test to be significant at a given significance level, denoted by α . Following the recommendations of the developers of this method for preliminary investigations, we can threshold on the Q value, such that we obtain an $FDR \leq \alpha$.

In linear models in which the transcript is the dependent variable and $\ln Lp-PLA_2$ is the covariate, we can use the P value on the beta-coefficient for $\ln Lp-PLA_2$ as a measure of the association of the transcript with $\ln Lp-PLA_2$. By the Q Value method, we observed 10 transcripts under the Q value threshold for $FDR \leq \alpha \leq .05$ (table 2 and fig. 4). To test for *cis*-regulation, which is defined as regulatory elements located at the gene,^{82–84} we computed the pointwise LOD score at the location reported in the NCBI and UniGene databases for each of the 341 transcripts within the 1-LOD interval of the $\ln Lp-PLA_2$ linkage signal. We take as our significance level a LOD score of 1.44, which is equivalent to $\alpha \leq .01$. Using the Q Value method for P values computed from the LOD scores, and thresholding for $FDR \leq \alpha \leq .01$, we observed seven transcripts (table 3 and fig. 4). Two of these—namely, the gene encoding Fc gamma receptor IIA (*FCGR2A* [MIM 146790]) (LOD 2.75) and the gene encoding heat-shock protein (70 kDa) (*HSP70*) (LOD 13.82) (*HSP70* = *HSPA6* [MIM 140555])—are presented in tables 2 and 3. We interpret these results to mean that these two genes—namely, *FCGR2A* and *HSP70*—are good candidates to pursue for further study.

FCGR2A is the receptor for C-reactive protein (CRP), a well-known acute-phase protein of the inflammation process involved in vascular dysfunction.^{85–87} Immunocyto-

Table 3. Transcripts under the 1-LOD Interval of the Linkage Signal on Chromosome 1 that Show *cis*-Regulation

Symbol	Name	Function	Location ^a (cM)	P ^b	Q
<i>ARNT</i>	Aryl hydrocarbon receptor nuclear translocator	Xenobiotic metabolism	147.72	8.46×10^{-5}	.00421
<i>MGC31963</i>	Chromosome 1 ORF 85	ORF	153.20	4.75×10^{-19}	0
<i>FCRH3</i>	Fc receptor-like protein 3	Immunoglobulin receptor	154.59	2.98×10^{-5}	.00178
<i>FCGR2A</i>	Fc gamma receptor 2A	C-reactive-protein receptor	158.42	.00019	.00693
<i>HSPA6 = HSP70</i>	Heat-shock protein (70 kDa)	Chaperone-protein folding	158.44	7.52×10^{-16}	0
<i>FCGR2B</i>	Fc fragment of immunoglobulin	Mast-cell activation	158.49	2.23×10^{-5}	.00167
<i>ATF6</i>	Activating transcription factor 6	Transcription factor	158.60	3.41×10^{-11}	0

^a Locations are averaged interpolations against our map, with use of physical distances obtained from the UCSC genome browser (Human [*Homo sapiens*] Genome Browser Gateway) for the two markers flanking the linkage peak at 153 cM.

^b P values are computed from the nominal LOD score at the location given in the NCBI and UniGene databases.

chemical work has shown that *FCGR2A* is highly expressed in the proliferative zones of atherosclerotic lesions.⁸⁸ Moreover, it has been shown in human monocytes that have the H131 mutation at the *FCGR2A* gene that *FCGR2A* has a significantly decreased binding to CRP⁸⁷ and that this mutation seems to confer protection against peripheral atherosclerosis.⁸⁹ The increased expression of heat-shock proteins, including HSP70, is known to be associated with an induced inflammatory response.⁹⁰⁻⁹³ Consistent with this knowledge, it has been noted that *HSP70* is overexpressed in several cell types, including monocytes, macrophages, and smooth-muscle cells, in advanced atherosclerotic lesions.⁹³

It should be noted that the beta-coefficients indicating the relationship of the transcript to rLp-PLA₂ were negative in sign for both transcripts (for *FCGR2A*, $\beta = -0.13$; for *HSP70*, $\beta = -0.12$). This is consistent with our observation of a decreasing QTL variance in rLp-PLA₂ from the nonobese to the obese-adiposity environment. To further characterize these transcripts, we performed multiple linear-regression analyses of the transcript expression levels as the independent variable and of age, sex, their second-order terms, and a relevant clinical trait as the dependent predictors. The analyzed clinical traits were diabetes status, impaired glucose tolerance, insulin resistance, adiposity status as defined above, dyslipidemia status, hypertension status, history of heart attack, history of heart surgery, and common and internal carotid artery intima-media thicknesses. For definitions of diabetes status, impaired glucose tolerance, insulin resistance, dyslipidemia status, and hypertension status, we again referred to the criteria suggested by the WHO.²⁸ There were significant associations between *FCGR2A*-expression levels and adiposity status ($P = .01636$), insulin resistance ($P = .00255$), and dyslipidemia status ($P = 1.3 \times 10^{-5}$). There was one clearly significant association between *HSP70*-expression levels and dyslipidemia status ($P = .02578$). The association between *HSP70*-expression levels and insulin resistance was barely nonsignificant ($P = .05613$). All other associations for both transcripts were nonsignificant. Taken together, the linkage and gene expression analyses indicate that we have identified a gene that is involved in

the inflammation process and is negatively regulated by the adiposity environment.

It is notable that, more than a decade ago, Després and colleagues proposed a hypothesis similar to the one addressed herein.⁹⁴⁻⁹⁶ It was suggested that visceral adiposity was capable of modulating the genetic susceptibility to coronary heart disease. Our results are supportive of that original hypothesis. In particular, we conclude that there is a significant polygenic and QTL genetic response to the adiposity environment in Lp-PLA₂, which is an important biomarker of inflammation and oxidative stress; that G \times E interaction analyses can improve our ability to identify and localize QTLs; and that there are at least two strong candidate genes underlying the QTL that we identified. Identification of genes and their variants involved in the inflammatory and oxidative processes in the metabolic syndrome is of high significance not only for the understanding of its metabolic pathogenesis but also for the development of therapeutic strategies to reduce the mortality and morbidity of this 21st-century epidemic.

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Web Resources

The URLs for data presented herein are as follows:

Human (*Homo sapiens*) Genome Browser Gateway, <http://genome.ucsc.edu/cgi-bin/hgGateway> (for physical distances)
NCBI, <http://www.ncbi.nlm.nih.gov/> (for the physical location of the transcripts)
Online Mendelian Inheritance in Man (OMIM), <http://www.ncbi.nlm.nih.gov/Omim/> (for *FCGR2A* and *HSPA6*)
Q Value, <http://faculty.washington.edu/~jstorey/qvalue/> (for a free software download)
SOLAR, <http://www.sfbr.org/solar/index.html> (for a free software download)

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